-27-WHAT IS CLAIMED IS: A method for recovering a desired target nucleic 1. acid molecule from a sample containing a mixture or library of single-stranded nucleic acid containing said molecule, wherein said method comprises the steps: 5 incubating said sample containing said nucleic Α. acid mixture or library in the presence of a primer nucleic acid molecule complementary to a sequence of said desired target molecule; said incubation being under conditions sufficient to 10 permit hybridization between said primer and said desired target molecule, and further sufficient to permit the template-dependent extension of said primer to thereby generate a double-stranded desired target molecule; 15 transforming single-stranded and double-stranded В. members of said mixture or library into a host cell, and recovering said desired molecule from said cell. C. The method of claim 1, wherein prior to 2. 20 commencing step A, said method comprises the presteps: incubating an initial sample containing said (1) nucleic acid mixture or library in the presence of a haptenylated nucleic acid probe molecule, said probe molecules having a sequence 25 complimentary to a nucleotide sequence of said desired target molecule; said incubation being under conditions sufficient to permit said probe to hybridize to said desired target molecule and to thereby generate a hybridized molecule 30 wherein said target molecule is bound to said probe;

-28incubating said sample containing said nucleic (2) acid mixture or library and biotinylated probetarget hybridized molecules of prestep (1) in the presence of a binding ligand of the hapten of said haptenylated probe, said binding ligand 5 being conjugated to support; said incubation being sufficient to permit said probe molecules, and said probe-target hybridized molecule to become bound to said binding ligand of said support; 10 recovering said probe-target hybridized (3) molecules bound to said support from said nucleic acid mixture or library and any unbound biotinylated probe-target hybridized molecules of prestep (2); and 15 incubating said recovered support containing (4) said bound probe-target hybridized molecules under conditions sufficient to separate the strands of double-stranded molecules; said incubation thereby releasing said hybridized 20 target molecule from said biotinylated probe, and generating a sample single-stranded desired target molecule for use in step (A). The method of claim 1, wherein said single-3. stranded nucleic acid molecule of said sample contains a 25 nucleotide analog, and wherein after completing step A, but prior to commencing step B, said method additionally comprises the presteps: (1') incubating said generated double-stranded molecules in the presence of a nuclease capable 30 of degrading nucleic acid containing nucleotide analog residues; and

- (2') incubating non-degraded nucleic acid with a primer under conditions sufficient to permit said primer to be extended in a templatedependent manner.
- 4. The method of claim 1, wherein in step A, said template-dependent extension of said primer is conducted in the presence of a nuclease resistant nucleotide analog to thereby generate a double-stranded desired target molecule containing a residue of said nucleotide analog; and wherein prior to commencing said step B, said method additionally comprises the presteps:

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- (1") incubating said generated double-stranded desired target molecule in the presence of a nuclease, wherein said nuclease is substantially unable to cleave a nucleic acid molecule containing said nucleotide analog residue, but is substantially capable of degrading both single-stranded nucleic acid molecules and double-stranded nucleic acid molecules that lack said nucleic acid amalog residue; said incubation being under conditions sufficient to permit such degradation, and thereby substantially eliminating both single-stranded nucleic acid molecules and double-stranded nucleic acid molecules that lack said nucleic acid analog residue from said sample; and thereby forming a preparation having a substantial enrichment of said desired target molecule relative to said initial sample; and (2") recovering said desired molecule from said
  - (2") recovering said desired molecule from said preparation of prestep (1") to thereby form a library or mixture for said step B.

- 5. The method of claim 1, wherein in step A said desired incubation is under conditions which minimize random hybridization.
- 6. The method of claim 2, wherein in prestep (1) 5 said desired incubation is under conditions which minimize random hybridization.
  - 7. The method of claim 1, wherein said desired target nucleic acid molecule is a DNA molecule.
- 8. The method of claim 7, wherein said DNA molecule is a single-stranded DNA molecule.
  - 9. The method of claim 1, wherein said desired target nucleic acid molecule îs an RNA molecule.
- 10. The method of claim 1, wherein said desired target nucleic acid molecule is a single-stranded nucleic acid molecule.
  - 11. The method of claim 1, wherein said desired target molecule is a circular nucleic acid molecule.
  - 12. The method of claim 11, wherein said desired target molecule is a circular DNA molecule.
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  13. The method of claim 2, wherein said hapten is biotin, and wherein said binding ligand of said hapten is avidin, streptavidin, or an antibody or antibody fragment that binds biotin.

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- 14. The method of claim 13, wherein said binding ligand of biotin is avidin.

Salin

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- 15. The method of claim 13, wherein said binding ligand of biotin is streptavidin.
- 16. The method of claim 2, wherein said support of said prestep (2) is a paramagnetic bead.
- 5 17. The method of claim 16, wherein said haptenylated probe-target hybridized molecule bound to said paramagnetic bead is recovered by magnetic means.
- 18. The method of claim 2, wherein in said primer molecule of step A is complementary to the same sequence of said desired target molecule as said probe molecule of substep (1).
  - 19. The method of claim 2, wherein in said primer molecule of step A is complementary to a sequence of said desired target molecule that differs from the sequence of said desired target molecule that is complementary to said probe molecule of substep (1).
  - 20. The method of claim 3, wherein said nucleic acid analog is deoxyuridine, and wherein said nuclease is UDG.
- 21. The method of claim 4, wherein said nuclease does not cleave hemimethylated DNA.
  - 22. The method of claim 21, wherein said nucleic acid analog is 5-methylcytidine, and wherein said nuclease that does not cleave hemimethylated DNA is HhaI.
- 23. The method of claim 2, wherein in prestep (1),
  25 said probe has a degenerate sequence.

- 24. The method of claim 4, wherein in step A, said primer has a degenerate sequence.
- 25. The method of claim 1, wherein said host cell is a bacterium.
- 26. The method of claim 1, wherein said method additionally includes the step of amplifying said desired target molecule by an in vitro amplification reaction.
  - 27. The method of claim 26, wherein said in vitro amplification reaction is a polymerase chain reaction.